

Package: peramo (via r-universe)

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Type Package

Title Permutation Tests for Randomization Model

Version 0.1.4

Description Perform permutation-based hypothesis testing for randomized experiments as suggested in Ludbrook & Dudley (1998) <[doi:10.2307/2685470](https://doi.org/10.2307/2685470)> and Ernst (2004) <[doi:10.1214/08834230400000396](https://doi.org/10.1214/08834230400000396)>, introduced in Pham et al. (2022) <[doi:10.1016/j.chemosphere.2022.136736](https://doi.org/10.1016/j.chemosphere.2022.136736)>.

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Suggests multcompView (>= 0.1-9)

BugReports <https://github.com/phamdn/peramo/issues>

License GPL-3

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peramo-package	<i>peramo: Permutation Tests for Randomization Model</i>
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Description

Perform permutation-based hypothesis testing for randomized experiments as suggested in Ludbrook & Dudley (1998) [doi:10.2307/2685470](https://doi.org/10.2307/2685470) and Ernst (2004) [doi:10.1214/088342304000000396](https://doi.org/10.1214/088342304000000396), introduced in Pham et al. (2022) [doi:10.1016/j.chemosphere.2022.136736](https://doi.org/10.1016/j.chemosphere.2022.136736).

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Author(s)

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AB *Two-Group Permutation Test*

Description

AB performs A/B testing for two-group experiments.

Usage

```
AB(a, b, rand = 9999, seed = 1)
```

Arguments

a	the measurement of responses of the first group.
b	the measurement of responses of the second group.
rand	an integer, the number of randomization samples. The default value is 9999.
seed	an integer, the seed for random number generation. Setting a seed ensures the reproducibility of the result. See set.seed for more details.

Value

AB returns a one-row data frame with 6 columns:

nA	the sample size of the first group.
mean.A	the mean responses of the first group.
nB	the sample size of the second group.
mean.B	the mean responses of the second group.
mean.dif	the difference between two mean responses.
pval	the p-value.

References

Ernst, M. D. (2004). Permutation Methods: A Basis for Exact Inference. *Statistical Science*, 19(4), 676–685. [doi:10.1214/088342304000000396](https://doi.org/10.1214/088342304000000396).

Examples

```
AB(c(19, 22, 25, 26), c(23, 33, 40))
```

copper

Biomarker Responses of the Ragworms to Copper and Warming

Description

Biomarker Responses of the Ragworms to Copper and Warming

Usage

Cu

bnk_worm

Format

An object of class data.frame with 60 rows and 6 columns.

An object of class data.frame with 210 rows and 22 columns.

copper_temp

Biomarker Responses of the Ragworms to Copper and Warming

Description

Biomarker Responses of the Ragworms to Copper and Warming

Usage

Cu

worm

Format

An object of class data.frame with 60 rows and 6 columns.

An object of class data.frame with 210 rows and 26 columns.

ctm	<i>Biomarker Responses of the Ragworms to Copper and Warming</i>
-----	--

Description

Biomarker Responses of the Ragworms to Copper and Warming

Usage

ctm_Cu

ctm_worm

Format

An object of class data.frame with 60 rows and 7 columns.

An object of class data.frame with 210 rows and 26 columns.

diffcalc	<i>Calculate the Differences between Means</i>
----------	--

Description

Calculate the Differences between Means

Usage

```
diffcalc(vec, control)
```

Arguments

vec a numeric vector, the mean responses.

control a logical, whether the control group exists.

ensulizole

Biomarker Responses of the Blue Mussels to Organic UV Filters

Description

Biomarker Responses of the Blue Mussels to Organic UV Filters

Usage

mussel_SoS

mussel_gill

mussel_digest

Format

An object of class `data.frame` with 15 rows and 7 columns.

An object of class `data.frame` with 120 rows and 24 columns.

An object of class `data.frame` with 120 rows and 24 columns.

Source

Pham, D. N., Sokolov, E. P., Falfushynska, H., & Sokolova, I. M. (2022). Gone with sunscreens: Responses of blue mussels (*Mytilus edulis*) to a wide concentration range of a UV filter ensulizole. *Chemosphere*, 309, 136736. doi:[10.1016/j.chemosphere.2022.136736](https://doi.org/10.1016/j.chemosphere.2022.136736).

nolesser

Compare the Differences with Critical Values

Description

Compare the Differences with Critical Values

Usage

```
nolesser(obs, cric)
```

Arguments

`obs` a numeric, the observed difference.

`cric` a numeric, the critical values of maximum absolute differences.

owl *One-Way Layout Permutation Test*

Description

owl performs the global test and multiple comparisons for single factor experiments.

Usage

```
owl(df, rand = 9999, alpha.post = 0.05, type.post = "control", seed = 1)
```

Arguments

df	a data frame with the name of experimental groups as the first column and the measurement of responses as the remaining columns.
rand	an integer, the number of randomization samples. The default value is 9999.
alpha.post	a numeric, the Type I error rate for multiple comparisons. The default value is 0.05.
type.post	the way of multiple comparisons, "all" for pairwise comparisons or "control" for only comparisons with the control group.
seed	an integer, the seed for random number generation. Setting a seed ensures the reproducibility of the result. See set.seed for more details.

Details

The first name appearing in the first column will determine the control group. The other names will be treatment groups.

Value

owl returns a list with 9 components:

n.obs	the sample sizes.
avg.obs	the mean responses.
T.obs	the T statistic for global test.
pval	the p-value for global test.
pval.round	the reported form of p-value.
main.test	the strength of evidence against the null hypothesis.
d.multi.obs	the differences in means for multiple comparisons.
mad.cric	the critical value of maximum absolute differences in means.
post.test	TRUE if the differences are significant.

References

Ernst, M. D. (2004). Permutation Methods: A Basis for Exact Inference. *Statistical Science*, 19(4), 676–685. doi:10.1214/088342304000000396.

Muff, S., Nilsen, E. B., O’Hara, R. B., & Nater, C. R. (2022). Rewriting results sections in the language of evidence. *Trends in Ecology & Evolution*, 37(3), 203–210. doi:10.1016/j.tree.2021.10.009.

Examples

```
ernst2004 <- data.frame(
  group = factor(rep(c("style1", "style2", "style3"), each = 5 ),
  levels = c("style1", "style2", "style3")),
  speed = c( 135,91,111,87, 122, 175,130,514,283, NA,105,147,159,107,194))
owl(ernst2004, type.post = "all")
```

owlStat

Test Statistics for One-Way Layout Permutation Test

Description

owlStat computes statistics for owl. This is not meant to be called directly.

Usage

```
owlStat(lov, env = parent.frame())
```

Arguments

lov	a list of vectors, responses by experimental groups.
env	an environment, to access outer scope variables.

Value

owlStat returns a list with 5 components:

n	the sample sizes.
avg	the mean responses.
T	the T statistic for global test.
d.multi	the differences in means for multiple comparisons.
mad	the maximum absolute differences in means.

.

References

Ernst, M. D. (2004). Permutation Methods: A Basis for Exact Inference. *Statistical Science*, 19(4), 676–685. doi:10.1214/088342304000000396.

twl	<i>Two-Way Layout Permutation Test</i>
-----	--

Description

twl performs the global test and multiple comparisons for two-factor experiments.

Usage

```
twl(  
  df,  
  rand = 4999,  
  seed = 1,  
  mult = FALSE,  
  simple = TRUE,  
  control = TRUE,  
  alpha = 0.05  
)
```

Arguments

df	a data frame with the first and second columns containing the levels of the two main factors and the third column containing the measurement of responses.
rand	an integer, the number of randomization samples. The default value is 4999.
seed	an integer, the seed for random number generation. Setting a seed ensures the reproducibility of the result. See set.seed for more details.
mult	a logical, whether to perform multiple comparisons.
simple	a logical, whether to perform comparisons for simple effects.
control	a logical, whether to perform only comparisons with the control group.
alpha	a numeric, the Type I error rate for multiple comparisons. The default value is 0.05.

Details

The first levels appearing in the first and second columns will determine the control groups (if any). The other levels will be treatment groups.

Value

twl returns a list with possible components:

n, n.main1, and n.main2	the sample sizes.
avg, avg.main1, and avg.main2	the mean responses.

`Fs` the F statistics, p-values, reported form of p-value, and strength of evidence against the null hypotheses.

`d.main1sub` and `d.main2sub` or `d.main1` and `d.main2`
the differences in means for multiple comparisons.

`mad.main1sub.cric` and `mad.main2sub.cric` or `mad.main1.cric` and `mad.main2.cric`
the critical value of maximum absolute differences in means.

`mult.test.main1sub` and `mult.test.main2sub` or `mult.test.main1` and `mult.test.main2`
TRUE if the differences are significant.

References

Manly, B. F. J. (2007). Randomization, bootstrap, and Monte Carlo methods in biology (3rd ed). Chapman & Hall/ CRC.

Ernst, M. D. (2004). Permutation Methods: A Basis for Exact Inference. *Statistical Science*, 19(4), 676–685. doi:10.1214/088342304000000396.

Muff, S., Nilsen, E. B., O'Hara, R. B., & Nater, C. R. (2022). Rewriting results sections in the language of evidence. *Trends in Ecology & Evolution*, 37(3), 203–210. doi:10.1016/j.tree.2021.10.009.

Motulsky, H. (2020). GraphPad Statistics Guide. GraphPad Software Inc. <https://www.graphpad.com/guides/prism/latest/statistics/index.htm>.

Examples

```
manly2007 <- data.frame(
  month = factor(rep(c("jun", "jul", "aug", "sep"), each = 6 ),
  levels = c("jun", "jul", "aug", "sep")),
  size = factor(rep(c("small", "large"), each = 3, times = 4),
  levels = c("small", "large")),
  consume = c( 13,242,105,182,21,7,8,59,20,24,312,68,515,488,88,460,1223,990,18,44,21,140,40,27))
twl(manly2007)
twl(manly2007, mult = TRUE, simple = TRUE, control = FALSE)
#might take more than 5s in some machines
```

twlStat

Test Statistics for Two-Way Layout Permutation Test

Description

twlStat computes statistics for twl. This is not meant to be called directly.

Usage

```
twlStat(df, env = parent.frame())
```

Arguments

df	a data frame with the levels of the two main factors as the first and second columns and the measurement of responses as the third column.
env	an environment, to access outer scope variables.

Value

twlStat returns a list with at least 4 components:

Fs	the F statistics for global test.
F.main1 and F.main2	the F statistics for the first main factor and the second main factor.
F.int	the F statistic for the interaction.

In case of multiple comparisons, additional components are:

avg or avg.main1 and avg.main2	the mean responses for multiple comparisons.
d.main1sub and d.main2sub or d.main1 and d.main2	the differences in means.
mad.main1sub and mad.main2sub or mad.main1 and mad.main2	the maximum absolute differences in means.

References

Manly, B. F. J. (2007). Randomization, bootstrap, and Monte Carlo methods in biology (3rd ed). Chapman & Hall/ CRC.

tw_complex

Permutation Test for Two-Way Layout with Extra Factors

Description

tw_complex performs the permutation test for ANOVA of two-factor experiments with complex design.

Usage

```
tw_complex(df, res, mains, nested, nuis, seed = 1, rand = 1999, emm = TRUE)
```

Arguments

df	a data frame with at least three columns.
res	a character string, name of response variable.
mains	two character strings, names of two main factors.
nested	(optional) a character string, name of the nested factor.
nuis	(optional) a character string, name of the nuisance factor.
seed	an integer, the seed for random number generation. Setting a seed ensures the reproducibility of the result. See set.seed for more details.
rand	an integer, the number of randomization samples. The default value is 1999.
emm	a logical, whether to compute estimated marginal means.

Details

res, mains, nested, and nuis refer to column names in df. While nuis column must be a numeric vector, mains and nested columns must be factors. res can be a numeric or logical vector.

tw_complex currently support linear models with only mains, generalized linear mixed-effects models with mains and nested, and linear mixed-effects models with mains, nested, and nuis.

Value

tw_complex returns a list with 3 main components:

lm, glmer, or lmer	model results.
anova	anova table.
perm	permutation test results with F-statistics, p-values, and strength of evidence.

References

Manly, B. F. J. (2007). Randomization, bootstrap, and Monte Carlo methods in biology (3rd ed). Chapman & Hall/ CRC.

Ernst, M. D. (2004). Permutation Methods: A Basis for Exact Inference. *Statistical Science*, 19(4), 676–685. doi:10.1214/088342304000000396.

Anderson, M., & Braak, C. T. (2003). Permutation tests for multi-factorial analysis of variance. *Journal of Statistical Computation and Simulation*, 73(2), 85–113. doi:10.1080/00949650215733.

See Also

[lm](#), [glmer](#), and [lmer](#).

Examples

```
tw_complex(df = subset(ctm_Cu, run == "Jan",
select = c("copper", "temp", "sediment")),
res = "sediment",
mains = c("copper", "temp"))
```

```
#might take more than 5s in some machines
```

XY

Permutation Test for Correlation Between Paired Samples

Description

XY performs permutation test on correlation coefficients.

Usage

```
XY(
  a,
  b,
  rand = 9999,
  seed = 1,
  use = "everything",
  method = c("pearson", "kendall", "spearman")
)
```

Arguments

a	a numeric vector, the first variable.
b	a numeric vector, the second variable.
rand	an integer, the number of randomization samples. The default value is 9999.
seed	an integer, the seed for random number generation. Setting a seed ensures the reproducibility of the result. See set.seed for more details.
method	correlation coefficient, "pearson", "kendall", or "spearman".

Value

XY returns an one-row data frame with 2 columns:

cor	the correlation coefficient.
pval	the p-value.

See Also

[cor.test](#)

Examples

```
with(subset(ctm_Cu, run == "Jan"), XY(sediment, porewater))
```

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